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The sequence and *de novo* **assembly of the giant panda genome**

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Abstract

Using next-generation sequencing technology alone, we have successfully generated and assembled a draft sequence of the giant panda genome. The assembled contigs (2.25 gigabases (Gb)) cover approximately 94% of the whole genome, and the remaining gaps (0.05 Gb) seem to contain carnivore-specific repeats and tandem repeats. Comparisons with the dog and human showed that the panda genome has a lower divergence rate. The assessment of panda genes potentially underlying some of its unique traits indicated that its bamboo diet might be more dependent on its gut microbiome than its own genetic composition. We also identified more than 2.7 million heterozygous single nucleotide polymorphisms in the diploid genome. Our data and analyses provide a foundation for promoting mammalian genetic research, and demonstrate the feasibility for using next-generation sequencing technologies for accurate, cost-effective and rapid *de novo* assembly of large eukaryotic genomes.

> The giant panda, *Ailuropoda melanoleura*, is at high risk of extinction because of human population expansion and destruction of its habitat. The latest molecular census of its population size, using faecal samples and nine microsatellite loci, provided an estimate of only 2,500–3,000 individuals, which were confined to several small mountain habitats in Western China¹. The giant panda has several unusual biological and behavioural traits, including a famously restricted diet, primarily made up of bamboo, and a very low fecundity rate. Moreover, the panda holds a unique place in evolution, and there has been continuing controversy about its phylogenetic position². At present, there is very little genetic

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Author Information This Whole Genome Shotgun project has been deposited at DDBJ/EMBL/GenBank under the accession ACTA00000000. The version described in this paper is the first version ACTA01000000. The reads data have been deposited in the EBI/NCBI Short Read Archive under the accession SRA009203. All panda data have been released for public use and can be freely accessed at BGI official website: [http://panda.genomics.org.cn/download.jsp.](http://panda.genomics.org.cn/download.jsp) The entire dataset includes all usable reads, genome assembly, annotation of genes, repeat and non-coding RNA, as well as heterozygous SNP, indel and structural variation information. Reprints and permissions information is available at www.nature.com/reprints. This paper is distributed under the terms of the Creative Commons Attribution-Non-Commercial-Share Alike licence, and is freely available to all readers at www.nature.com/nature.

information for the panda, which is an essential tool for detailed understanding of the biology of this organism.

A major limitation in obtaining extensive genetic data is the prohibitive costs associated with sequencing and assembling large eukaryotic genomes. The development of nextgeneration massively parallel sequencing technologies, including the Roche/454 Genome Sequencer FLX Instrument, the ABI SOLiD System, and the Illumina Genome Analyser, has significantly improved sequencing throughput, reduced costs, and advanced research in many areas, including large-scale resequencing of human genomes^{3,4}, transcriptome sequencing, messenger RNA and microRNA expression profiling, and DNA methylation studies. However, the read length of these sequencing technologies, which is much shorter than that of traditional capillary Sanger sequencing reads, has prevented its use as the sole sequencing technology in *de novo* assembly of large eukaryotic genomes.

Here, using only Illumina Genome Analyser sequencing technology, we have generated and assembled a draft genome sequence for the giant panda with an assembled N50 contig size (defined in Table 1) reaching 40 kilobases (kb), and an N50 scaffold size of 1.3 megabases (Mb). This represents the first, to our knowledge, fully sequenced genome of the family Ursidae and the second of the order Carnivora⁵. We also carried out several analyses using the complete sequence data, including genome content, evolutionary analyses, and investigation of some of the genetic features underlying the panda's unique biology. The work presented here should aid in understanding and carrying out further research on the genetic basis of panda's biology, and contribute to disease control and conservation efforts for this endangered species. Furthermore, our demonstration that next-generation sequencing technology can allow accurate *de novo* assembly of the giant panda genome will have farreaching implications for promoting the construction of reference sequences for other animal and plant genomes in an efficient and cost-effective way.

Short-read *de novo* **sequencing and assembly**

For sequencing, we selected a 3-year-old female giant panda from the Chengdu breeding centre in China. The panda genome contains 20 pairs of autosomes and one pair of sex chromosomes $(2n = 542)$ (Supplementary Fig. 1). We used a whole-genome shotgun sequencing strategy and Illumina Genome Analyser sequencing technology. DNA was extracted from the peripheral venous blood, and 37 paired-end sequencing libraries were constructed with insert sizes of about 150 base pairs (bp), 500 bp, 2 kb, 5 kb and 10 kb. In total, we generated 176-Gb of usable sequence (equal to 73-fold coverage of the whole genome), with an average read length of 52 bp (Supplementary Tables 1 and 2).

We assembled the short reads using SOAPdenovo [\(http://soap.genomics.org.cn](http://soap.genomics.org.cn))—a genome assembler developed specifically for use with next-generation short-read sequences⁶ (Supplementary Fig. 2). SOAPdenovo uses the de Bruijn graph algorithm⁷ and applies a stepwise strategy to make it feasible to assemble the panda genome using a supercomputer (32 cores and 512 Gb random access memory (RAM)). The algorithm is sensitive to sequencing errors, so we excluded the data generated from poor libraries, filtered lowquality reads, and used the 134 Gb (56-fold coverage) high-quality reads for *de novo* assembly.

We first assembled the short reads from fragmented small insert-size libraries (<500 bp) into contigs using sequence overlap information. Contigs were not extended into regions in which repeat sequences created ambiguous connections. At this point, we assembled about 39-fold coverage short-reads into contigs having an N50 length of 1.5 kb, achieving a total length of 2.0 Gb (Table 1). Here, we avoided using reads from long insert-size paired-end

libraries (2 kb) on contig assembly because these libraries were constructed using a circularization and random fragmentation method⁴, and the small fraction $(\sim 5\%)$ of chimaeric reads in these long insert-size libraries could generate incorrect sequence overlap resulting in misassembly.

We then used the paired-end information, step by step from the shortest (150 bp) to the longest (10 kb) insert size, to join the contigs into scaffolds. We obtained a scaffold N50 length of 1.3Mband a total length of 2.3 Gb, determined by counting the estimated intrascaffold gaps. Most of the remaining gaps probably occur in repetitive regions, so we further gathered the paired-end reads with one end mapped on the unique contig and the other end located in the gap region and performed local assembly with the unmapped end to fill in the small gaps within the scaffolds. The resulting assembly had a final contig N50 length of 40 kb (Table 1). In total, 223.7 Mb gaps were closed. Roughly 54.2Mb (2.4% of total scaffold sequence) remained unclosed, of which we determined that about 90% contained carnivorespecific transposable elements and the remainder were primarily tandem repeats with high unit identity and lengths larger than the sequencing read length, which could not be assembled with the current data. About 0.05% of the panda assembly was composed of tandem repeats (Supplementary Table 3). Given the genome similarity between the panda and the dog, and that 0.2% of the dog genome is made up of tandem repeats, we estimate that around 3.6Mb (0.15%) of tandem repeat sequences might be missing in the current panda genome assembly.

To evaluate the single-base accuracy of the assembled genome sequence, we realigned all the usable sequencing reads onto the scaffolds using SOAPaligner⁸. The peak sequencing depth was 65×, and more than 20 reads covered over 99% of the assembled sequences (Fig. 1a). Using massively parallel next-generation sequencing technology and improving sequencing depth to more than $20 \times$ coverage provides a very high single-base accuracy^{3,9}.

The GC content difference is a primary factor for non-random sequencing-depth distribution⁴. Using 500-bp non-overlapping sliding windows along the genome, which is similar to the size of sequencing DNA fragments (150–500 bp), we found that both very low and very high GC regions had a relatively lower sequencing depth, but nearly all regions with a GC content between 20% and 80% had more than 20x coverage, which is sufficient for *de novo* assembly (Supplementary Fig. 3). Only a minor fraction (0.004%) of the panda genome—as compared to the sequenced dog (0.079%), human (0.095%) and mouse (0.015%) genomes—had a GC content lower than 20% or higher than 80% (Supplementary Fig. 4). Assuming similar GC content in the panda and dog genomes, 0.075% (1.8 Mb) sequence with extreme GC content could be missing in the current panda assembly. Nevertheless, the analysis showed that *de novo* genome assembly is unlikely to be strongly affected by GC-biased non-random sampling.

There are 27 known panda mRNA genes in GenBank, one of which is the SRY sex determination gene located on chromosome Y, thus not present in the female panda. We were able to detect the remaining 26 genes in the assembled scaffolds with 99.3% total sequence aligned (Supplementary Table 4). All of these genes, with the exception of the ribosomal gene *RPS15*, were intact on the scaffolds. For *RPS15*, we found one fragment in the scaffolds and the other in a small contig not incorporated in the scaffolds. These data indicate that the genome assembly has good coverage and completeness for genes with unique sequences, but given their nature, multiple-copy genes or genes containing repetitive sequences may be fragmented.

To assess the large-scale and local assembly accuracy of the scaffolds, we sequenced and assembled nine bacterial artificial chromosomes (BACs) independently using Sanger

sequencing technology¹⁰. Each BAC (average length 97.7 kb) was aligned to only one scaffold, and up to 98.0% of the total BAC regions were well covered by the assembled contigs. We did not observe any obvious misassembly errors (Fig. 1b and Supplementary Fig. 5). We then checked the single-base mismatch or small insertion/deletion differences. After excluding the annotated heterozygous single nucleotide polymorphisms (SNPs) and indels (described later), the rate of single-base differences was 0.07% and the average number of insertions/deletions on each BAC was 16 (Supplementary Table 5 and Supplementary Fig. 6). There was both a very high read depth on the scaffolds and a high Phred score on the BACs at the discrepant bases, so most of these might be unidentified heterozygous SNPs. Manual inspection showed that 38% of the insertions/deletions were caused by assembly errors on the BAC (Supplementary Fig. 7a), and the rest were unidentified heterozygous indels of the diploid genome (Supplementary Fig. 7b).

To assess genome coverage of the assembled contigs and scaffolds, we first estimated the genome size of the panda. To obtain the best estimate, we used sequencing depth, size ratio of syntenic blocks, and comparison of C-values. Using 17-base oligonucleotides and their frequency distribution in the usable sequencing reads to determine sequencing depth¹¹, we obtained an estimated panda genome size of 2.46 Gb (Supplementary Fig. 8). Given the presence of sequencing errors, we expect that the sequencing depth is underestimated, and consequently the panda genome size should be slightly smaller than 2.46 Gb.

Given the evolutionary relationship of the panda with the dog and the bear, we used their genome size information to refine our estimate of the panda genome size. For the whole genome, the size ratio of conserved syntenic blocks between the panda and the dog genomes is 0.978. As the current estimated size of the dog genome is 2.45 Gb^5 , we estimated the panda genome size to be 2.40 Gb. Genome sequence and size information for the bear is unavailable. We therefore used C-values (haploid DNA content in picograms), as this is proportional to genome size. The recorded C-value is 2.80 for the dog and 2.75 for two bears, *Ursus arctos* and *Ursus hibetanus* [\(http://www.genomesize.com/\)](http://www.genomesize.com/), indicating that the genome size of bears should be smaller than that of the dog. On the basis of previous reports showing that the karyotype of panda is similar to that of bears¹², the bear and the panda are expected to have similar genome sizes. We again determined the panda genome size to be 2.40 Gb. Using this genome size, the assembled contigs and scaffolds should cover about 94% and 96% of the whole genome, respectively.

Panda, dog and human repeat comparison

Combining results from analyses using the Repbase library¹³, alignment with known transposable-element-related genes, and *de novo* RepeatModeller¹⁴ identification, we estimate that transposable elements comprised approximately 36.2% of the panda genome, which is similar to that of the dog genome (36.1%), and lower than the human genome (46.1%) (see Supplementary Tables 6 and 7). Analysis of the divergence rate of the transposable element in the panda genome using Repbase transposable-element libraries showed that nearly all of the identified panda transposable-element copies had $a > 10\%$ divergence rate from the consensus. This high divergence rate may be related to the fact that the Repbase transposable-element consensus sequences were annotated using mammalian genomes other than the panda. Using RepeatModeller transposable elements, we found that about 70Mb of transposable-element sequences (3% of the genome) had $a < 10\%$ divergence rate from the consensus (Supplementary Fig. 9), which are likely to be active transposable elements of recent origin. (See Supplementary Results for further analyses on specific transposable elements.)

Panda genome has a low divergence rate

To investigate the rate of recent segmental duplication $(>90\%$ identity, >1 kb length) in the panda genome, we conducted self-sequence alignment and identified 3,095 duplicated fragments with a total length 10.4 Mb (0.43%) in the whole genome assembly. To determine whether these highly similar recent duplicates were collapsed by the assembly, we aligned all sequencing reads to the assembled genome, and identified 5,485 segments (>1 kb in length) with a total length of 13.9 Mb that had an obviously higher read depth than the other genomic regions. Considering the average depth was about 2.47 times that of the whole genome, we estimated that the total length of the duplicated copies was about 34.3 Mb (1.43% of the whole genome) (Supplementary Table 8). Integrating the results of the two methods, we estimated that recent segmental duplication comprised about 43.7Mb (1.82%) of the panda genome, which is lower than the rate in the dog genome $(106.6 \text{ Mb}, 4.21\%)^{15}$.

To investigate panda genome conservation and evolution, we performed whole-genome alignment with the panda, dog and human genomes. Each of these genomes contained \sim 1.4Gb of non-repetitive sequence, of which 846 Mb of sequence was shared by all three genomes. Of the remainder, 322 Mb of sequence was shared between panda and dog, which was much higher than that between panda and human (163 Mb) or between dog and human (58 Mb). The panda had the lowest amount of lineage-specific sequences, indicating that it has the lowest divergence rate among the three genomes (Fig. 2a).

We found that the panda, dog and human genomes had high genomic synteny. For example, the syntenic relationship of 35 panda scaffolds, 36.5Mb of human chromosome 2, and dog chromosome37 (Fig. 2b) showed no large-scale rearrangement within the chromosomal region. For the whole genome, regions of conserved synteny between the panda and dog genomes covered 2.22 Gb (96.7%) and 2.27 Gb (92.9%), respectively; and the pairwise syntenic regions between the panda and human genomes covered 2.19 Gb (95.3%) and 2.66 Gb (88.1%), respectively (Supplementary Table 9).

We further detected genomic rearrangement events using the whole genome alignments. Here, we used 3,805 panda scaffolds with lengths longer than 10 kb, which covered 98.6% of the assembled genome. To define rearrangements, we used different cutoffs for minimum syntenic segments. The total intra-chromosomal breakpoints between the panda and the dog genomes varied from 468 with a 5-kb cutoff to 20 with a 100-kb cutoff. We also identified about 42 inter-chromosomal rearrangements using a 100-kb cutoff (Supplementary Table 10). We used the human genome as the common outgroup and found that there were 4–5 times more rearrangements in dog than in panda, which provided evidence that the panda has a lower divergence rate than the dog.

Gene prediction and comparison

To predict the number of genes in the panda genome, we used both evidence-based and *ab inito* methods. We aligned 20,001 (94%) of the human and 19,086 (99%) of the dog genes to the panda genome and predicted 19,303 and 19,245 gene loci, respectively (see Methods and Supplementary Table 11). We also used Genscan¹⁶ and Augustus¹⁷ with model parameters trained on the human, and predicted 44,428 and 29,238 gene loci, respectively for the two programs, in the panda genome. To facilitate further analysis, we integrated all the gene sources and created a reference gene set that contained 21,001 genes for the panda (Supplementary Table 12).

To measure the quality of gene prediction, we compared the length distribution of genes, coding sequences (CDS), exons and introns, and the distribution of exon number per gene among representative mammalian genomes, including panda, dog, human, mouse and

opossum. The panda was similar to the human with respect to all of these key parameters (Supplementary Fig. 10). To evaluate the rate of missing exons in the predicted genes, we aligned both the panda and dog genes to the human genes and calculated the percentage of human gene sequences covered. On average, the dog and panda genes covered 96.2% and 93.5% of the human gene sequences, respectively (Supplementary Fig. 11a). Further checking showed that most (more than 80%) of the unannotated exons were at the 5′ or 3′ ends of genes; these exons were usually very small and separated by large introns that were hard to predict by gene finders, although they were correctly presented in the assembled scaffolds (Supplementary Fig. 11b). There are 626 genes, with predicted functions in defence/ immunity, receptors and signalling, located in the segmental duplication regions, which was similar to the findings in a previous analysis of the dog genome¹⁵ (Supplementary Table 13). Overall, we found that the quality of the predicted panda genes was comparable to that of other well-annotated mammalian genomes.

We next used InParanoid and Multiparanoid¹⁸ to identify gene orthologues among panda, dog, mouse and human. In total, we found 18,643 orthologous clusters that were shared by at least two species, and 15,060 clusters that were shared by all four species (Supplementary Fig. 12). There were 2,534 panda-specific genes, which is nearly double the amount of dogspecific genes (1,677). This difference might be due to the fact that only an evidence-based method was used for dog gene prediction and a smaller number of genes was predicted in the dog genome $(19,305)^5$.

Gene gains and losses are one of the primary contributors to functional changes¹⁹. To obtain greater insight into the evolutionary dynamics of the genes, we determined the expansion and contraction of the gene orthologue clusters among these four species. We used a maximum-likelihood method to estimate the orthologue cluster sizes in their common ancestor 20 , and then defined the expansion and contraction by comparing the cluster size differences between the ancestor and each of the current species. Overall in all four genomes, we found a greater amount of gene contraction than expansion (Fig. 3), indicating that loss-of-function might have an important role in functional evolution. We also saw that the clusters within different species that underwent the most marked changes (expansion or contraction) were those involved in receptor activity (Supplementary Table 14).

We looked for signatures of positive selection using branch-site likelihood ratio tests²¹ for 12,424 genes of high confidence that had 1:1 orthologues in the panda and human genome and in at least one of the rat, mouse and dog genomes that had passed a series of rigorous filters for large-scale synteny, alignment quality, conservation of exon-intron structure, and sequence quality²². Using three different tests— one specific for the panda lineage, one specific for the dog lineage, and one combining evidence from all five species included in the alignment—we found 134, 94 and 182 positively selected genes (PSGs), respectively, using a conservative 5% false-discovery-rate criterion.

The panda and the dog lineage only share six PSGs. However, Fisher's exact and Mann-Whitney *U* tests showed enrichment of several immunity- and defence-related functional categories in both the panda and the dog genomes, which is consistent with the results from previous genome-wide positive selection scans in mammalian genomes²³ (Supplementary Table 15). Gene Ontology classification showed that panda PSGs spanned a diverse range of immune response systems, in particular acute inflammatory and innate immune response, as well as response to wounding (Supplementary Table 16).

Of note, the inclusion of the panda genome in the multiple species analysis showed an enrichment of two new Gene Ontology categories 'blood circulation and gas exchange activity' and 'haemostasis' as compared to an analysis with six mammalian species²³

(human, chimpanzee, macaque, mouse, rat and dog). In contrast to the various enrichments for Gene Ontology categories for the panda PSGs, only two categories, 'cytolysis' and 'cellcell adhesions', were over-represented in the dog PSGs. The latter category includes cadherins (for example, *PCDHB10* (*P*=2.66×10−32), *PCDHB8* (*P*=1.80×10−19), *PCDHB17* (*P*=3.74×10−10) and *ME1* (*P*=7.10×10−9)).

Panda-specific characteristics

To gain insight into some of the traits unique to the panda, we investigated genes that may influence characteristics such as panda diet and fecundity. For our investigation of genes that may be involved in food selection and digestion, we identified genes that encode digestive enzymes protease, amylase, lipase, cellulase, lactase, invertase and maltase in the panda genome, indicating that the panda probably has all the necessary components for a carnivorous digestive system. We did not find any homologues of digestive cellulase genes, including endoglucanase, exoglucanase and beta-glucosidase, indicating that the bamboo diet of the panda is unlikely to be dictated by its own genetic composition, and may instead be more dependent on its gut microbiome.

Taste is also an important factor in the development of dietary habits. The five components of basic taste are: sweetness, saltiness, sourness, bitterness and umami. Bitterness is sensed by the *T2R* gene family24, and we identified 16 *T2R* genes in the panda genome, which is comparable to 14 *T2R* genes reported for the dog²⁵. Umami is sensed through the *T1R* gene family26. In the panda genome, *T1R2* and *T1R3* are in an intact form, but *T1R1* has become a pseudogene (Fig. 4)—we found that the two panda *T1R1* exons contain frameshift errors, which we confirmed by Sanger sequencing. We also found that the d_N/d_S (the ratio of the rate of non-synonymous substitutions to the rate of synonymous substitutions) difference for this gene was low between the panda (0.17) and dog (0.13) lineage, indicating a recent death for this gene in the panda lineage (Supplementary Fig. 13). The T1R1/T1R3 heterodimer is known to be a receptor that senses umami²⁷, and a previous study indicated that $TIR1/T1R3$ may be the sole receptor for umami taste²⁸. Umami perception occurs through the detection of the carboxylate anion of glutamic acid, which is a naturally occurring amino acid common in meats, cheese, broth, stock and other protein-heavy foods. Thus, the loss-offunction of the *T1R1* gene might prevent the panda from expressing a functional umami taste receptor, which may partly explain why the panda diet is primarily herbivorous despite its taxonomic classification as a carnivore.

The giant panda has a low fecundity rate, which is of great consequence given its endangered status. We identified nearly all of the reproduction genes known to be critical for mammalian gonad development and function. We identified a putative pseudo folliclestimulating hormone (FSH) β-subunit gene (giant panda-*FSHB2*), which has high sequence similarity to the giant panda FSH β-subunit gene (giant panda-*FSHB1*) in the panda genome as well as in several other mammalian species, including cow and horse (Supplementary Fig. 14). At this stage, whether the pseudo *FSHB2* gene contributes to the reproductive features of the giant panda remains to be determined.

Panda has a high heterozygosity rate

As the panda population is estimated to be \sim 2,500 wordwide at present¹, the issue of genetic variability and evolutionary potential is extremely important because low genetic variability will cause a population decline. A recent molecular analysis of the 655 bp of the mitochondrial control region DNA and ten microsatellite loci indicated that the panda still has high genetic variability²⁹. Sequence data for the whole genome to firmly establish this important issue have not been available until now.

Using the assembled panda genome sequence as a reference, we realigned all of the usable sequencing reads with the genome to identify the heterozygous SNPs, small indels, and structural variations between the two sets of homologous chromosomes in the dip-loid genome. We were able to align about 85% of the usable sequencing reads. Among these, we used the 90% that had unique best locations to detect variation by applying the pipeline developed in the analysis of the first Asian human genome^{3,6,8} (Supplementary Table 17). We identified 2.7 million heterozygous SNPs in the panda diploid genome. The estimated heterozygosity rate of the autosomes was 1.35×10^{-3} for the whole genome and 0.66×10^{-3} for the coding regions (Fig. 5a). This is about 1.95 times higher than the rate estimated for the human genome $(0.6\times10^{-3}$ and $0.34\times10^{-3})^3$. The heterozygosity rate in the X chromosome is about half that of the autosomes, and the ratio of transition/transversion is 2.1 (Supplementary Table 18). Our finding of a high heterozygosity rate may be due to the mixed genetic background (Liangshan and Minshan sub-populations) of the panda that we sequenced. These results, however, may indicate that the panda species still has high genetic variability, which would support the potential for its successful survival despite its small population size. To determine this, analyses of other panda genomes are required.

The heterozygous SNP rate in the panda genome showed a bimodal distribution with one peak at 0.004% and another at 0.035% (Fig. 5b). The SNP rate of 14.9% of the genome regions was lower than 0.015%, and the SNP rate of the remaining regions was in range 0.015–0.6%. According to the Wright-Fisher population genetic model³⁰, the effective population size is inversely proportional to the time to the most recent common ancestor (TMRCA), and TMRCA directly determines the heterozygosity, so this SNP distribution might indicate changes or bottlenecks of the effective population size in the panda population history.

In addition to SNPs, we identified 267,958 small indels that ranged in size from 1 to 6 bp (Supplementary Table 19). The estimated small indel rate was 1.22×10^{-4} and 0.70×10^{-4} on autosomes and sex chromosomes, respectively—roughly one order of magnitude lower than that of the SNP rate. We also identified insertions or deletions that were greater than 100 bp and inversion events. There were 4,359 insertions and deletions detected with a median length of 150 bp, and 20 inversions (Supplementary Table 20). More than 70% of the insertions or deletions overlapped transposable elements, and these were primarily made up of carnivore-specific LINE/L1 and SINE/Lys transposable elements. We expect that the amount of structural variation was underestimated because the genome assembly is still too fragmented to identify long structural variations.

Discussion

Comparative genomics is a powerful tool for determining the genetic basis of biological functions. Sequencing more mammalian genomes will undoubtedly facilitate our ability to annotate the human genome. So far, however, only a few mammals have had their complete genomes sequenced, and a great deal more data are necessary for carrying out more detailed mammalian comparative genomics. The primary limitations are related to the extreme cost and length of time needed to sequence and assemble very large genomes that would be suitable for such analyses.

Our ability to generate and assemble a draft sequence for an entire mammalian genome using only next-generation Illumina Genome Analyser short-read sequencing technology indicates that such technology can be used to generate many more mammalian genome draft sequences in a rapid and cost-effective manner. The assembled contigs and scaffolds of the panda genome were sufficiently long to allow us to carry out our gene prediction and comparative analyses, and our evaluation of the sequence quality showed that the panda

genome sequence was of similar utility to sequences generated using traditional capillarybased Sanger sequencing. It should be noted that fewer active transposons and recent segmental duplications in the panda genome may have made it relatively easier to assemble than some other complex animal or plant genomes. Thus, the relative affect of these genomic features on *de novo* assembly from short reads will need to be assessed in the future in genomes that contain such features.

The panda genome is, to our knowledge, the first genome from the Ursidae lineage, and the second carnivore to be sequenced. Our comparative analysis showed that the panda had high genomic synteny with dog and human, but that there has been considerable divergence in the repetitive regions, most of which seems to result from recent transposable-element activity. Our comparison of the panda and other available mammalian genomes showed that most genes were conserved among all of these mammals, although we also identified several gene families that had expanded or contracted within each lineage. Of interest, our analysis of genes potentially involved in the evolution of the panda's reliance on bamboo in its diet showed that the panda seems to have maintained the genetic requirements for being purely carnivorous even though its diet is primarily herbivorous. Furthermore, given our finding that some of the genes necessary for complete digestion of bamboo are missing from its genome, investigation of panda's gut microbiome may be important for understanding its unusual dietary restrictions.

As well as providing a reference genome for use in future studies, we were also able to detect the most extensive heterozygous SNPs, indels and structural variation in the diploid genome, which will be important for ecological and population studies designed to address issues of panda survival.

METHODS SUMMARY

We constructed the paired-end DNA libraries with insert sizes larger than 2 kb by selfligation of the DNA fragments and merging the two ends of the DNA fragment. We randomly fragmented the circularized DNA and enriched the fragments crossing the merged boundaries using magnetic beads with biotin and streptavidin. The sequencing process followed the manufacturer's instructions (Illumina), and the fluorescent images were processed to sequences using the Illumina data processing pipeline (v1.1).

The genome sequence was assembled with short reads using SOAPdenovo software⁶ ([http://](http://soap.genomics.org.cn) [soap.genomics.org.cn\)](http://soap.genomics.org.cn), which adopts the de Bruijn graph data structure to construct contigs⁷ . The reads were then realigned to the contig sequence, and the paired-end relationship between the reads was transferred to linkage between contigs. We constructed scaffolds starting with short paired-ends and then iterated the scaffolding process, step by step, using longer insert size paired-ends. To fill the intra-scaffold gaps, we used the pairedend information to retrieve read pairs that had one read well-aligned on the contigs and another read located in the gap region, then did a local assembly for the collected reads.

Known transposable elements were identified using RepeatMasker (version $3.2.6$)¹⁴ against the Repbase³¹ transposable element library (version 2008-08-01), and highly diverged transposable elements were identified with RepeatProteinMask¹⁴ by aligning the genome sequence to the curated transposable-element-related proteins. A *de novo* panda repeat library was constructed using RepeatModeller¹⁴. Using evidence-based gene prediction, the human and dog genes (Ensembl release 52) were projected onto the panda genome, and the gene loci were defined by using both sequence similarity and whole-genome synteny information. *De novo* gene prediction was performed using Genscan¹⁶ and Augustus¹⁷. A reference gene set was created by merging all of the gene sets. The sequencing reads were

mapped on the panda genome sequence using SOAPaligner⁸, and heterozygous SNPs were identified by SOAPsnp⁹.

Supplementary Material

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 $\mathbf a$

Percentage of genome

8

 $\,$ 6 $\,$

 $\overline{4}$

 $\overline{2}$

Figure 1. Assessment of assembly quality

a, Distribution of sequencing depth of the assembled genome. **b**, Comparison of the assembled genome with a BAC sequence. Read depth on the BAC was calculated by mapping the Illumina Genome Analyser short reads onto the BAC sequence. The predicted gene and annotated transposable elements (TEs) on the BAC sequence are shown in green and red, respectively. The remaining unclosed gaps on the scaffolds are marked as white blocks.

Figure 2. Conserved sequences among the panda, dog and human genomes

a, The total lengths of aligned and unaligned non-repetitive sequences. Each of the three genomes contains 1.4Gb of non-repetitive sequences. Pairwise whole-genome alignment was performed using Blastz. The lengths shown are in Mb. **b**, Syntenic view of the dog chromosome 37, 35 panda scaffolds (>10 kb), and the human chromosome 2. Note that the 3Mb region at the end of dog chromosome 37 was unanchored.

Figure 3. Dynamic evolution of orthologous gene clusters

The estimated numbers of orthologue groups in the common ancestral species are shown on the internal nodes. The numbers of orthologous groups that expanded or contracted in each lineage after speciation are shown on the corresponding branch, with '+' referring to expansion and '−' referring to contraction.

Figure 4. Structure of the umami receptor *T1R1* **gene**

Two frameshift mutations occurred in the third and sixth exons (red) of the panda *T1R1* gene. The third exon contained a 2-bp ('GG') insertion; the sixth exon contained a 4-bp ('GTGT') deletion.

a

Figure 5. Panda heterozygous SNP density

a, Statistics of identified heterozygous SNPs. Analysed regions are the genomic regions with proper unique read coverage that were used for heterozygote detection. The panda Xchromosome-derived scaffolds were identified by Blastz alignment to the dog X chromosome. **b**, Distribution of heterozygosity density in the panda diploid genome. Heterozygous SNPs between the two sets of chromosomes of the panda diploid genome were annotated, then non-overlapping 50-kb windows were chosen and heterozygosity density in each window was calculated.

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Table 1

Summary of the panda genome sequencing and assembly Summary of the panda genome sequencing and assembly

*** Approximate average insert size of Illumina Genome Analyser sequencing libraries. The sizes were estimated by mapping the reads onto the assembled genome sequences. High-quality read sequences that were used in assembly. Coverage was estimated assuming a genome size of 2.4 Gb. Sequence coverage refers to the total length of generated reads, and physical coverage $\frac{c}{c}$ *†*High-quality read sequences that were used in assembly. Coverage was estimated assuming a genome size of 2.4 Gb. Sequence coverage refers to the total length of generated reads, and physical coverage refers to the total length of sequenced clones of the libraries. refers to the total length of sequenced clones of the libraries. $*$ N50 size of contigs or scaffolds was calculated by ordering all sequences then adding the lengths from longest to shortest until the summed length exceeded 50% of the total length of all sequences. N90 is *‡*N50 size of contigs or scaffolds was calculated by ordering all sequences then adding the lengths from longest to shortest until the summed length exceeded 50% of the total length of all sequences. N90 is similarly defined. similarly defined.